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Sicegar: R package for sigmoidal and double-sigmoidal curve fitting

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ABSTRACT

Sigmoidal and double-sigmoidal dynamics are commonly observed in many areas of biology. Here we present *sicegar*, an R package for the automated fitting and classification of sigmoidal and double-sigmoidal data. The package categorizes data into one of three categories, “no signal”, “sigmoidal”, or “double sigmoidal”, by rigorously fitting a series of mathematical models to the data. The data is labeled as “ambiguous” if neither the sigmoidal nor double-sigmoidal model fit the data well. In addition to performing the classification, the package also reports a wealth of metrics as well as biologically meaningful parameters describing the sigmoidal or double-sigmoidal curves. In extensive simulations, we find that the package performs well, can recover the original dynamics even under fairly high noise levels, and will typically classify curves as “ambiguous” rather than misclassifying them. The package is available on CRAN and comes with extensive documentation and usage examples.

INTRODUCTION

Growth patterns resembling sigmoidal or double sigmoidal curves are common in biological systems. In ecological systems the number of individuals in a population may follow a sigmoidal growth pattern over time, while in molecular systems the production of a reporter may have similar dynamics. Quantifying and comparing these kinds of growth dynamics can lead to valuable insight into the systems’ behavior. However, the automated quantification and classification of thousands of noisy growth curves, in a high-throughput manner, can be technically challenging.

Here we present the *sicegar* package, an R package designed to automatically analyze and classify time course growth data. The input data is assumed to represent intensity measured over time, where intensity represents any metric of a system that may vary in a sigmoidal or double-sigmoidal pattern. Using these time course data, *sicegar* categorizes the data as “sigmoidal”, “double sigmoidal”, “ambiguous”, or “no signal”, by first checking the data for the presence of signal (i.e., the intensity curve deviates significantly from a horizontal line and varies significantly) and then fitting two mathematical models to the data, a sigmoidal one and a double-sigmoidal one. The package automatically determines which curve provides the better fit, and it reports a wide range of parameters about the fitted models that enable easy comparison of the fitted curves within and among experiments.

The *sicegar* package was originally written to study poliovirus infection and replication at the single-cell level (Guo et al., 2017), and the package name is inspired by this application (SINGLE CELL Growth Analysis in R). However, the potential applications for the package go beyond this one use, and the package is designed as a generalized way to classify growth data from time course experiments. In the following, we describe the mathematical models used to classify the data, how the models are fit, and the inferences that can be made from them.

42 METHODS

43 Model fitting and categorization

44 We implemented R code that can fit data to the sigmoidal and double-sigmoidal curves defined by Eq. 1
45 and Eq. 5 in Results and Discussion. Before fitting, data is internally normalized to a $[0, 1]$ interval on
46 both the x and y axis. The sigmoidal and double sigmoidal models are then fit to the internally normalized
47 data via likelihood maximization, using the function `nls.lm` in the `minpack.lm` package (Elzhov
48 et al., 2015). To guarantee robust fitting `nls.lm` is run multiple times using different starting estimates,
49 chosen at random from a uniform distribution covering the allowed interval of parameters. For each fit,
50 the quality of fit based on AIC scores is recorded, and the parameter estimates that produce the overall
51 best fit across replicates are reported as the final estimates.

52 The simplest way to use `sicegar` is via the function `fitAndCategorize`, which attempts to
53 both identify the best-fitting model and determine whether this model is sigmoidal, double-sigmoidal, or
54 neither. This function first checks whether there is any signal in the data, by testing whether the maximum
55 observed intensity exceeds a user-defined threshold and the data range also exceeds a user-defined
56 threshold. If either test fails, the data are labeled as “no signal”.

57 Next, the function fits both a sigmoidal and a double-sigmoidal curve to the data and then assess which
58 provides the better fit. This process considers a number of different factors. First, it assesses whether both
59 models could actually be fitted (the fitting did not fail). Second, the AIC scores of the fitted models are
60 examined to make sure they are greater than a threshold value (the default is -10). The starting time points
61 of each model are then checked to insure that they are positive. The starting intensities are examined to
62 insure that they are smaller than a threshold (the default is 0.05). Then the ratio of the models’ intensity
63 prediction at the last observation to the models’ maximum intensity prediction is tested to insure that it is
64 over 0.85 in the case of the sigmoidal model and below 0.75 in the case of the double sigmoidal model.
65 If any of these tests fail for either the sigmoidal or double-sigmoidal fit, then the data is not categorized
66 as being described by that model. If both models fail any of these tests the data is categorized as being
67 “ambiguous”. If only one model passes these tests while the other fails, then the data is considered to
68 be best described by that model. If both of the fits pass all of these test a decision is made about which
69 model best describes the data based on their AIC scores. If the sigmoidal AIC score is smaller than the
70 double sigmoidal AIC score then the data is labeled as “sigmoidal”, if it is larger then the data is labeled
71 as “double sigmoidal”.

72 The complete decision process, as well as the specific parameters users can modify to adjust the
73 package to their own needs, are explained in detail in the vignette “Identifying the best-fitting model
74 category” supplied with the package.

75 Simulations

76 To assess `sicegar` performance, we simulated sigmoidal and double-sigmoidal data with different levels
77 of noise. We first generated 50 distinct sets of parameters for both sigmoidal and double-sigmoidal curves.
78 For the sigmoidal model, the maximum, slope, and midpoint parameters were chosen uniformly at random
79 from the intervals $[0.3, 20]$, $[0.001, 40]$, and $[3, 27]$, respectively. For the double-sigmoidal model, the
80 maximum, increasing slope, the first midpoint, the decreasing slope, the distance between midpoints, and
81 final asymptote intensity ratio parameters were chosen uniformly at random from the intervals $[0.3, 20]$,
82 $[0.001, 40]$, $[3, 26]$, $[0.001, 40]$, $[1, 27 - \text{first midpoint}]$, and $[0, 0.85]$, respectively. Using these parameters
83 we simulated data from $t = 3$ to $t = 30$ in increments of 0.5, for a total of 55 time points each.

84 For each parameter set we then generated samples that included noise. We used 11 distinct noise
85 amplitudes, ranging from 0% to 150% of the maximum value of the initial simulated data. We generated
86 three noise vectors with the same maximum amplitude for each noise level and create a unique noise
87 vector for each replicate. To generate a noise vector at a given amplitude for a sample, we created a vector
88 of random numbers chosen from the uniform distribution between -0.5 and 0.5 for each time value in the
89 dataset. We then multiplied each random number with the noise amplitude and the maximum intensity of
90 the dataset. Finally, we added these noise values to the intensity values of the initial samples and obtained
91 noisy samples. In total, this procedure generated 1650 sigmoidal and double-sigmoidal noisy samples
92 with varying levels of uniform noise, for a total of 3300 samples.

93 For each simulated sample, we used `sicegar` to estimate both the shape of the curve (“sigmoidal”,
94 “double-sigmoidal”, or “ambiguous”) and the curve’s original parameter values. To estimate the algo-
95 rithm’s accuracy in fitting a model to the data, we calculated the mean absolute error between the initial

96 (without noise) and predicted intensity vectors, and we normalize this number relative to the maximum
 97 intensity of the initial sample. We calculated the normalized mean absolute error regardless of whether
 98 the algorithm predicted the category of the sample correctly or not.

99 **Package availability and documentation**

100 The `sicegar` package is implemented in R and available from CRAN at
 101 <https://CRAN.R-project.org/package=sicegar>. The `sicegar` source code is available
 102 on github at <https://github.com/wilkelab/sicegar>. Extensive user documentation is avail-
 103 able in multiple vignettes distributed as part of the package.

104 The package depends on existing R packages `dplyr` (Wickham and Francois, 2015), `minpack.lm`
 105 (Elzhov et al., 2015), `fBasics` (Team et al., 2014), and `ggplot2` (Wickham, 2009).

106 **RESULTS AND DISCUSSION**

107 Growth in many biologically systems can be thought of as occurring in two phases (Fig. 1). In the first
 108 phase, intensity will increase exponentially until saturation is reached and it levels off at some asymptotic
 109 maximum level. In the second phase, the intensity may then decay from the maximum value to a lower
 110 one or even back to zero. Depending on the system and/or the length of the observation this second phase
 111 may or may not occur. These two phases can be modeled with two sigmoidal curves that describe the
 112 relationship between time and intensity. The first describes the increase of intensity and the second its
 113 decay.

114 Based on this two-phase view of growth an observer of a system may observe three different types of
 115 behaviors of intensity over time. First, the observer may observe the full process and clearly see both the
 116 growth and the decay phases, i.e., the full double-sigmoidal curve. Second, if the observation ends before
 117 decay occurs, or if the decay phase is not relevant for the system under study, the observer may only see
 118 the first sigmoidal curve. Third, if the intensity is not produced in detectable amounts, the observer will
 119 not see any signal at all. We label these three types of behavior as “double sigmoidal”, “sigmoidal”, and
 120 “no signal”. If a given time course does not clearly fit into any of these three categories we label it as
 121 “ambiguous”.

122 On the face of it, fitting a sigmoidal curve to time-course data appears to be a simple and solved
 123 problem. A sigmoidal curve is fully determined by only three parameters, and many non-linear curve-
 124 fitting algorithms exist that can be used to fit a simple three-parameter function. However, when we
 125 attempted to fit thousands of curves to experimentally measured data, we found that achieving numerical
 126 reliability was non-trivial. While a simple algorithm will work most of the time, it will fail eventually
 127 on cases that by manual inspection have a clear solution. The problem was further exacerbated for
 128 double-sigmoidal curves, where even guaranteeing that the curve is double-sigmoidal at all times was
 129 non-trivial (see Appendix). In the following, we describe the mathematical models and fitting procedures
 130 we developed to reliably obtain good fits and to be able to automatically categorize time courses into the
 131 different types of behavior.

132 **Sigmoidal and double-sigmoidal models**

A sigmoidal curve can be uniquely determined by three parameters, the maximum value, the midpoint,
 and the slope of the curve (Fig. 2A). To represent a sigmoidal curve, the `sicegar` package uses the
 Fermi function

$$I(t) = f_{\text{sig}}(t) = \frac{I_{\text{max}}}{1 + \exp(-a_1(t - t_{\text{mid}}))}. \quad (1)$$

133 Here, $I(t)$ is the intensity time course, given as a function of time t . The three parameters to be fitted
 134 are I_{max} , t_{mid} , and a_1 . The parameter I_{max} represents the maximum intensity observed, the parameter t_{mid}
 135 indicates the time at which intensity has reached half of its maximum, and the parameter a_1 is related to
 136 the slope of $I(t)$ at $t = t_{\text{mid}}$ via the formula $d/dt I(t)|_{t=t_{\text{mid}}} = a_1 I_{\text{max}}/4$ (Fig. 2A).

137 A double sigmoidal curve can be uniquely determined by six parameters, two midpoints and slopes
 138 parameters, a maximum value, and an asymptotic final value after decay (Fig. 2B). We define the double-
 139 sigmoidal model by combining two regular sigmoidal functions. This combined function rises from 0 to a
 140 maximum value I_{max} at time t^* and then decays to the final value I_{final} . The function is divided into two
 141 parts, one to the left of the maximum, for $t < t^*$ (the growth phase) and one to the right of the maximum,

142 for $t > t^*$ (the decay phase). The two parts are rescaled separately, such that the maximum value of $I(t)$ is
 143 given by I_{\max} and the final value of $I(t)$ for large times is given by $I_{\text{final}} > 0$.

To define our double-sigmoidal model, we begin with a simple double-sigmoidal function, obtained by multiplying together two sigmoidal functions:

$$f_{\text{dsig-base}}(t) = \frac{1}{1 + \exp(-a'_1(t - t'_{\text{mid1}}))} \frac{1}{1 + \exp(-a'_2(t - t'_{\text{mid2}}))}. \quad (2)$$

We then define the time t^* at which this function is maximal,

$$t^* = \operatorname{argmax} f_{\text{dsig-base}}(t), \quad (3)$$

and the value of the function at that time point,

$$f_{\max} = f_{\text{dsig-base}}(t^*). \quad (4)$$

We then write

$$I(t) = f_{\text{dsig}}(t) = \begin{cases} c_1 f_{\text{dsig-base}}(t) & \text{for } t \leq t^* \text{ (growth phase)} \\ c_2 f_{\text{dsig-base}}(t) + I_{\text{final}} & \text{for } t > t^* \text{ (decay phase)} \end{cases}, \quad (5)$$

with

$$c_1 = \frac{I_{\max}}{f_{\max}} \quad (6)$$

and

$$c_2 = \frac{(I_{\max} - I_{\text{final}})}{f_{\max}}. \quad (7)$$

144 This definition may seem awkward and overly complex, but it guarantees that certain non-sigmoidal
 145 corner cases are eliminated (see Appendix).

146 The meaning of the parameters in the double-sigmoidal model mirror those of the regular sigmoidal
 147 model. The time point t'_{mid1} determines (but does not exactly represent) the time at which the intensity
 148 has risen to half of its maximum, and the time point t'_{mid2} (enforced to be larger than t'_{mid1}) determines
 149 (but does not exactly represent) the time at which the intensity has decayed halfway from its maximum to
 150 its final value. The prime in t'_{mid1} and t'_{mid2} indicates that these time points are not the final parameters
 151 of interest. The final parameters of interest, the true midpoints t_{mid1} and t_{mid2} , have to be calculated
 152 numerically. Similarly, the parameters a'_1 and a'_2 determine the speed of growth and speed of decay,
 153 respectively, but do not represent the slopes at times t_{mid1} and t_{mid2} . The true midpoint slopes also have to
 154 be calculated numerically, by numerically differentiating f_{dsig} at each midpoint time. As stated before, the
 155 parameters I_{\max} and I_{final} represent the maximum intensity and the final intensity after decay, respectively.

156 In addition to the parameters that uniquely determine the sigmoidal and double-sigmoidal functions,
 157 the `sicegar` package calculates other useful metrics that describe aspects of the curves (Fig. 2). For the
 158 sigmoidal model, we calculate the starting time of growth by finding the point that a line which passes
 159 through the midpoint t_{mid} with the slope a_1 intersects with the x-axis. This value gives the length of time
 160 between the start of observations and the first observable increase in the response variable. Using the same
 161 line which passes through t_{mid} with slope a_1 , we can also determine the amount of time that has elapsed
 162 during exponential growth, by looking at the time point where the line intersects with the maximum
 163 intensity (I_{\max}) and subtracting the starting time. For the double-sigmoidal function the time until decay
 164 can be calculated in a similar way (Fig. 2B).

165 Performance against simulated data

166 To evaluate `sicegar`'s performance more systematically, we created simulated datasets with sigmoidal
 167 and double-sigmoidal curves to which we applied increasingly higher amounts of random noise, and we
 168 assessed how well `sicegar` performed on these datasets.

169 First, we studied how well `sicegar` recovers the type of the curve (sigmoidal or double-sigmoidal)
 170 at varying noise levels. For curves that were simulated as sigmoidal, `sicegar` virtually never identified

171 them as double-sigmoidal (Fig. 3A). However, starting at noise levels of around 45% (corresponding
172 to a noise amplitude of half the total amplitude of the sigmoidal curve), `sicegar` classified a fraction
173 of the curves as ambiguous. The fraction of curves classified as ambiguous increased with increasing
174 noise level until virtually all curves were classified as ambiguous at noise levels of around 100%. Results
175 were similar for curves simulated as double-sigmoidal, except that a small fraction of curves was always
176 identified as sigmoidal, regardless of noise level. These were generally curves for which the parameter
177 values happened to cause very little decay at large times, either because the final asymptotic value was
178 close to the maximum or because decay started late relative to the maximum time point simulated. Finally,
179 at high noise levels, double-sigmoidal curves can appear sigmoidal, and `sicegar` classified those curves
180 as such.

181 We also compared the fits `sicegar` obtained from noisy data to the initial curves before applying
182 noise. For each fit, we calculated the normalized mean absolute error, which is the mean of the absolute
183 differences of fitted and original intensity levels, normalized by the maximum original intensity. We found
184 that these errors were near zero for low noise levels and increased only gradually as noise increased (Fig
185 3B). Thus, overall, our algorithm results in reliable fits, it fails gradually with increasing noise levels, and
186 it is conservative in assessing whether it has correctly identified a sigmoidal or double-sigmoidal curve or
187 not.

188 Comparison to existing packages

189 The `sicegar` package provides a generalized way to analyze growth dynamics. However, `sicegar` is
190 not the only package with the ability to fit data to these types of models. The `qPCR` (Ritz and Spiess, 2008)
191 package fits linear, exponential, a number of sigmoidal, and log-logistic functions to qPCR (realtime
192 quantitative polymerase chain reaction) data. Unlike `sicegar` this package was designed to analyze
193 a specific type of data (qPCR) and uses a log-logistic function rather than a double sigmoidal function
194 to describe data that includes both a growth and decay phase. A similar package, `grofit` (Kahm
195 et al., 2010), uses a number of parametric growth curves and a model-free spline approach to estimate
196 parameters from dose-response data. This package does not include a parametric model that would
197 account for a decay phase in the data, though an interface to construct custom models is provided. The
198 intended use of this package is to model pharma-/toxicological data and it produces estimates of values
199 such as half maximum effective concentration (EC50). While `sicegar` can estimate these same types
200 of parameters, it was designed for more general usage. For example, the first midpoint estimated by
201 `sicegar` could refer to the EC50, though depending on the system it may or may not make sense to refer
202 to that point in those terms. Another package, `FlexParamCurve` (Oswald et al., 2012), fits in excess
203 of 30 models to data. While this package is useful for teasing apart exactly which variant of a model
204 best describe dynamics, `sicegar` is better suited for classifying data as either containing sigmoidal or
205 double-sigmoidal dynamics.

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APPENDIX: PROPERTIES AND DERIVATIONS OF THE SIGMOIDAL AND DOUBLE-SIGMOIDAL MODELS

The `sicegar` package uses two distinct models that represent growth and growth-like data obtained from time course experiments. The sigmoidal model describes situations in which the signal starts from zero and rises to a maximum level, and the double-sigmoidal model describes situations in which signal starts from zero, rises to a maximum level, and then declines towards an asymptotic value below the maximum.

Two factors were considered when choosing the corresponding mathematical functions. First, the models should be as simple as possible while being well-behaved. Well-behaved implies that the models should not take on unexpected shapes under specific parameter combinations, they should have few variables, they should be continuous, and they should be defined for all time points from minus infinity to plus infinity. As we will see below, achieving these constraints is non-trivial for the double-sigmoidal model. Second, as much as possible, model parameters should have simple interpretations and so represent or be related to biologically meaningful parameters.

Properties of the sigmoidal function

The `sicegar` package uses for its sigmoidal fit the Fermi function (Eq. 1). Here we describe the properties of a more general variant of this function,

$$I(t) = \frac{I_{\max} - I_{\text{init}}}{1 + \exp[-a_1(t - t_{\text{mid}})]} + I_{\text{init}}. \quad (8)$$

The Fermi function is recovered by setting $I_{\text{init}} = 0$.

The more general function discussed here has four parameters, each of which has a clear meaning in terms of the shape of the function:

1. I_{init} represents the initial value of the sigmoidal curve at minus infinity,

$$I_{\text{init}} = \lim_{t \rightarrow -\infty} I(t). \quad (9)$$

2. I_{\max} represents the maximum value of the curve at plus infinity,

$$I_{\max} = \lim_{t \rightarrow \infty} I(t). \quad (10)$$

3. t_{mid} represents the time point at which the slope of $I(t)$ is maximal,

$$t_{\text{mid}} = \operatorname{argmax} \frac{d}{dt} I(t). \quad (11)$$

4. The parameter a_1 , constrained to be positive at all times, is related to the maximum slope of $I(t)$,

$$\frac{a_1}{4}(I_{\max} - I_{\text{init}}) = \left. \frac{d}{dt} I(t) \right|_{t=t_{\text{mid}}} \quad (12)$$

In the following, we will also need the inverse sigmoidal function,

$$I(t) = \frac{I_{\max} - I_{\text{final}}}{1 + \exp[a_2(t - t_{\text{mid}})]} + I_{\text{final}}, \quad (13)$$

which differs from Eq. 8 in that it does not have a minus sign in front of the slope parameter a_2 . We also have renamed I_{init} as I_{final} , to indicate that it now corresponds to the limit $t \rightarrow \infty$. The inverse sigmoidal function represents sigmoidal decay rather than sigmoidal growth, starting at its maximum and decaying to its final value.

Properties of the double-sigmoidal function

The double-sigmoidal function is defined in Eq. 5 in the main text. It rises from zero to a maximum value and then decays towards an asymptotic final value, and it is defined for all t . The function is continuous with a continuous first derivative and a discontinuous second derivative.

The parameters I_{\max} and I_{final} represent the maximum and asymptotic values of the curve, respectively. However, the parameters t'_{mid1} , t'_{mid2} , a'_1 , and a'_2 do not directly represent midpoints and slopes. Instead, the two midpoints and two slopes need to be calculated numerically.

Derivation of the double-sigmoidal function

The double-sigmoidal as defined in Eq. 5 may seem overly complex and awkward. However, we had to define it in this way to avoid problems that arise with other formulations.

In particular, a seemingly more straightforward double-sigmoidal could be obtained by multiplying together a regular sigmoidal (Eq. 8) and an inverse sigmoidal (Eq. 13):

$$I(t) = \left(\frac{I_{\max} - I_{\text{init}}}{e^{-a_1(t-t_{\text{mid1}})} + 1} + I_{\text{init}} \right) \left(\frac{1 - I_{\text{final}}}{e^{a_2(t-t_{\text{mid2}})} + 1} + I_{\text{final}} \right) \quad (14)$$

Even though this function behaves appropriately for most parameter choices, there are parameter regions in which this function is not a double sigmoidal. This occurs in general when the interval between t_{mid1} and t_{mid2} is very small, and one of the slopes is very steep while the other one is not. As an example, consider the case shown in Fig. 4, where Eq. 14 generates a local minimum in addition to a local maximum. These kinds of corner cases will be discovered by the fitting algorithm when used on sufficiently many diverse datasets, and therefore we ruled out Eq. 14 as an appropriate choice for our double-sigmoidal function.

However, Eq. 14 will never have a local minimum if we set both I_{init} and I_{final} to zero (see next section for proof). Thus, we define the *base double sigmoidal* as

$$f_{\text{dsig-base}}(t) = \frac{1}{(e^{-a_1(t-t_{\text{mid1}})} + 1)(e^{a_2(t-t_{\text{mid2}})} + 1)} \quad (15)$$

This function has the correct shape in principle, except that the intensity always decays back to zero for large t .

One possible solution to the problem that the function approaches zero for $t \rightarrow \infty$ is to cut it into two parts, rescale them separately, and then merge them together again. We cut the function at its maximum, where $t = t^*$. Thus, the part from $t = -\infty$ to $t = t^*$ represents sigmoidal growth and the part from $t = t^*$ to $t = \infty$ represents sigmoidal decay. Both parts are rescaled such that the desired I_{\max} and I_{final} are obtained. This procedure leads to the definition of the double-sigmoidal we use, as defined in Eq. 5. This function has the desired limiting properties and is guaranteed to have only one maximum. However, the cutting and rescaling procedure causes its second derivative to be discontinuous at $t = t^*$.

The base double sigmoidal has exactly one local maximum

For the base double sigmoidal as defined in Eq. 15 we can prove the following Lemma.

Lemma. $f_{\text{dsig-base}}(t)$ has one local maximum.

Proof. At its local maximums and minimums, i.e. when $t = t^*$, the derivative of $f_{\text{dsig-base}}(t)$ must be equal to zero,

$$\left. \frac{d}{dt} f_{\text{dsig-base}}(t) \right|_{t=t^*} = 0. \quad (16)$$

By definition, $t_{\text{mid2}} > t_{\text{mid1}}$, so without loss of generality we can write $t_{\text{mid2}} = t_{\text{mid1}} + L$, where L is a positive number. Then we obtain

$$\left. \frac{d}{dt} \left[\frac{1}{(e^{-a_1(t-t_{\text{mid1}})} + 1)(e^{a_2(t-L-t_{\text{mid1}})} + 1)} \right] \right|_{t=t^*} = 0 \quad (17)$$

We define new variables $u \equiv t - t_{\text{mid}1}$ and $u^* \equiv t^* - t_{\text{mid}1}$. We also define the function $g(u) = f_{\text{dsig-base}}(u + t_{\text{mid}1})$. Finding the roots of $g(u)$ is equivalent to finding the roots of $f_{\text{dsig-base}}(t)$, and it involves solving the following equation for u^* :

$$\left. \frac{d}{du} g(u) \right|_{u=u^*} = \left. \frac{d}{du} \left[\frac{1}{(e^{-a_1 u} + 1)(e^{a_2(u-L)} + 1)} \right] \right|_{u=u^*} = 0 \quad (18)$$

After taking the derivative, we can rewrite this equation as

$$\frac{e^{(a_1 u^* + a_2(L-u^*))} \left[a_1(e^{a_2(L-u^*)} + 1) - a_2(e^{a_1 u^*} + 1) \right]}{(e^{a_1 u^*} + 1)^2 (e^{a_2(L-u^*)} + 1)^2} = 0 \quad (19)$$

For the left-hand side to be equal to zero, the term in the square brackets in the numerator must be equal to zero, because all other terms in the expression are always positive. We next define the term in the square brackets as a function of u ,

$$h(u) = a_1(e^{a_2(L-u)} + 1) - a_2(e^{a_1 u} + 1). \quad (20)$$

The number of roots of $h(u)$ is equal to the number of local extrema of $f_{\text{dsig-base}}(t)$.

To determine the number of roots of $h(u)$, we first note the following limiting properties:

$$\lim_{u \rightarrow -\infty} h(u) = \infty \quad (21)$$

$$\lim_{u \rightarrow \infty} h(u) = -\infty \quad (22)$$

According to the mean value theorem, $h(u)$ must thus have at least one root. Next, we note that $h(u)$ is strictly decreasing:

$$\frac{d}{du} h(u) = -a_1 a_2 e^{a_2(L-u)} - a_1 a_2 e^{a_1 u} < 0 \quad (23)$$

Since the derivative of $h(u)$ is not equal to zero for any u we can use Rolle's Theorem to conclude that the function cannot have more than one root. In conclusion, $h(u)$ has exactly one root, which implies that $f_{\text{dsig-base}}(u)$ has exactly one local extremum.

Finally, we demonstrate that the local extremum of $f_{\text{dsig-base}}(t)$ is a local maximum, by inspecting the second derivative of $g(u)$ at $u = u^*$.

$$\begin{aligned} \left. \frac{d^2}{du^2} g(u) \right|_{u=u^*} &= \frac{e^{a_1 u - 2a_2(L-u)}}{(e^{a_1 u} + 1)^3 (e^{-a_2(L-u)} + 1)^3} \\ &\times \left[\left(a_1(e^{a_2(L-u)} + 1) - a_2(e^{a_1 u} + 1) \right)^2 \right. \\ &\quad \left. - a_1^2 e^{a_1 u} (e^{a_2(L-u)} + 1)^2 - a_2^2 (e^{a_1 u} + 1)^2 e^{a_2(L-u)} \right] \Big|_{u=u^*} \quad (24) \end{aligned}$$

The first term in the square brackets is equal to $h(u)^2$ and thus is zero at $u = u^*$. The two remaining terms inside the square brackets are strictly negative, and the term outside the square brackets is strictly positive. Therefore, $g(u^*) < 0$, which proves that the local extremum is a local maximum. \square

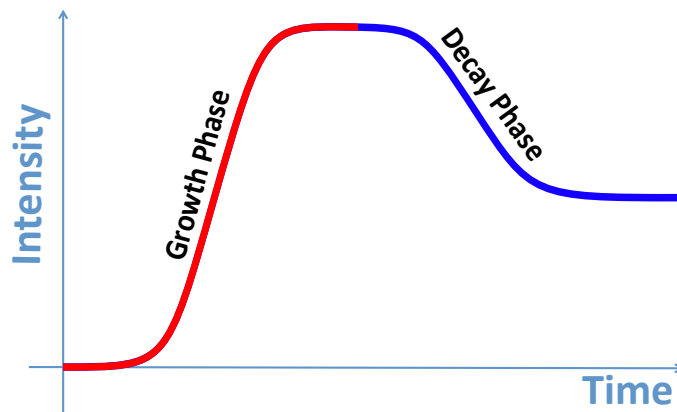


Figure 1. The two phases that an observer of growth data might see. The first phase, shown in red, is the “growth phase”, during which growth occurs at exponential rate until some asymptotic level is reached. The second phase, shown in blue, is the “decay phase”, during which intensity declines towards some lower asymptotic level.

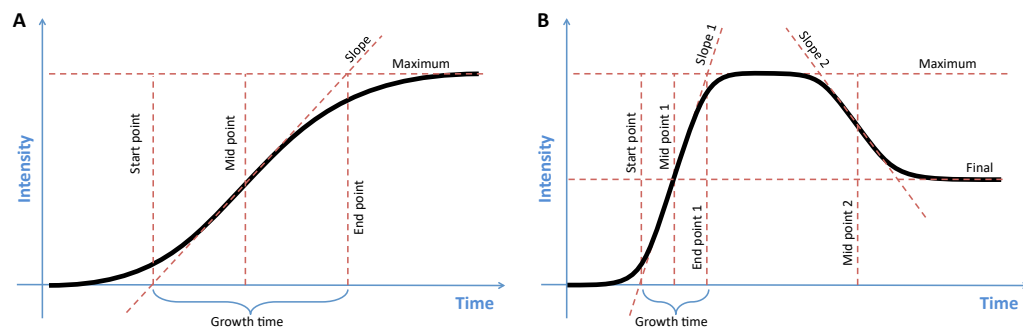


Figure 2. The two models used to classify growth data, with key parameters of each model labeled. (A) In the sigmoidal model the maximum value, the midpoint, and the slope of the curve uniquely describe the function. Estimates of the start and end points of growth, as well as the growth time, can then be calculated from this model. (B) The double sigmoidal model is uniquely determined by six parameters, two midpoint and two slope parameters, a maximum value, and a final value. As in the case of the sigmoidal model, the start point, end point, and duration of both the growth and the decay phases can be calculated from these parameters.

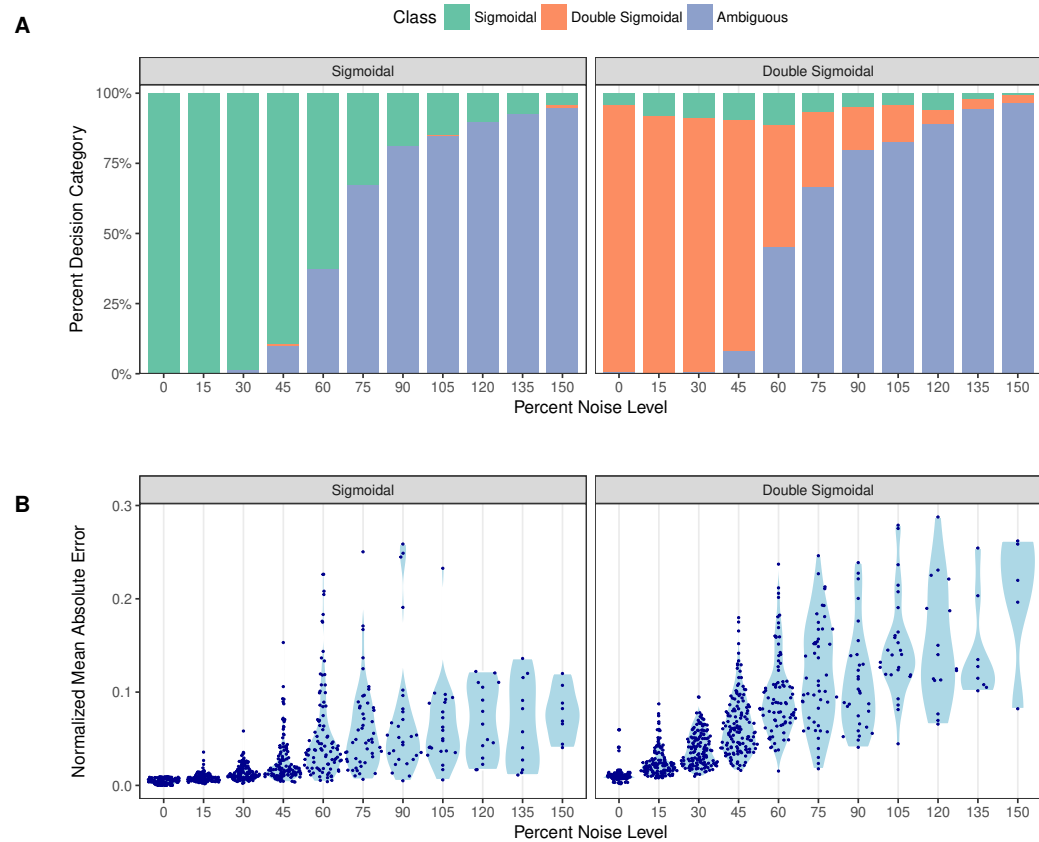


Figure 3. Assessment of *sicegar* performance on simulated data subjected to varying levels of noise. Noise levels are measured relative to the maximum intensity of the simulated curves. (A) *sicegar* tends to either recover the original type of curve correctly (for small to moderate amounts of noise) or label the dataset as ambiguous (at high noise levels). (B) The difference between the original data (without noise) and the fitted curve is near zero for low noise levels and increases gradually and slowly for higher noise levels.

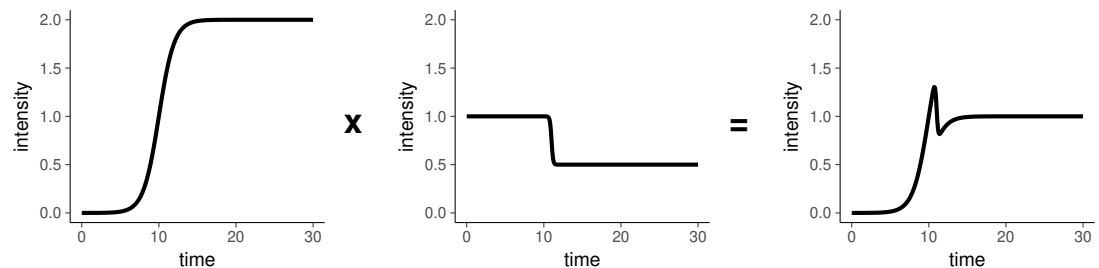


Figure 4. Example combination of parameters that are problematic for the multiplicative double-sigmoidal model (Eq. 14), which is obtained by multiplying the sigmoidal (Eq. 8) and inverse sigmoidal (Eq. 13) functions. Parameter choices for this example are: $I_{\text{init}} = 0$, $I_{\text{final}} = 0.5$, $I_{\text{max}} = 2$, $a_1 = 1$, $a_2 = 10$, $t_{\text{mid}1} = 10$, $t_{\text{mid}2} = 11$. This example demonstrates that Eq. 14 is not guaranteed to be double-sigmoidal.